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## Inference in mixed models in R - beyond the usual asymptotic likelihood ratio test

Slides

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# Inference in mixed models in R - beyond the usual asymptotic likelihood ratio test

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# History

- ▶ Years ago, Ulrich Halekoh and SH colleagues at “Danish Institute for Agricultural Sciences”
- ▶ That was SAS-country back then
- ▶ Many studies called for random effects models - and for `PROC MIXED`
- ▶ `PROC MIXED` reports (by default)  $p$ -values from asymptotic likelihood ratio test.
- ▶ Main concern: Effects should be “tested against” the correct variance component in order not to make effects appear more significant than they really are.

# History

- ▶ Common advice: Use Satterthwaite or Kenward-Roger approximation of denominator degrees of freedom in  $F$ -test – in an attempt not to get things “too wrong”.
- ▶ Then  $\mathbb{R}$  came along; we advocated the use of  $\mathbb{R}$ .
- ▶ Random effects models were fitted with the **nlme** package – but there was no Satterthwaite or Kenward-Roger approximation, so our common advice fell apart.

# History

## The degree of freedom police...

R-help - 2006: [R] how calculation degrees freedom

<https://stat.ethz.ch/pipermail/r-help/2006-January/087013.html>

SH: Along similar lines ... probably in recognition of the degree of freedom problem. It could be nice, however, if `anova()` produced ...

Doug Bates: I don't think the "degrees of freedom police" would find that to be a suitable compromise. :-)

In reply to another question:

Doug Bates: I will defer to any of the "degrees of freedom police" who post to this list to give you an explanation of why there should be different degrees of freedom.

# History

## Motivation: Sugar beets - A split-plot experiment

- ▶ Model how sugar percentage in sugar beets depends on harvest time and sowing time.
- ▶ Five sowing times ( $s$ ) and two harvesting times ( $h$ ).
- ▶ Experiment was laid out in three blocks ( $b$ ).

Experimental plan for sugar beets experiment

Sowing times:

1: 4/4, 2: 12/4, 3: 21/4, 4: 29/4, 5: 18/5

Harvest times:

1: 2/10, 2: 21/10

Plot allocation:

	Block 1					Block 2					Block 3					
	+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					
Plot	h1	h1	h1	h1	h1	h2	h2	h2	h2	h2	h1	h1	h1	h1	h1	Harvest time
1-15	s3	s4	s5	s2	s1	s3	s2	s4	s5	s1	s5	s2	s3	s4	s1	Sowing time
	+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					
Plot	h2	h2	h2	h2	h2	h1	h1	h1	h1	h1	h2	h2	h2	h2	h2	Harvest time
16-30	s2	s1	s5	s4	s3	s4	s1	s3	s2	s5	s1	s4	s3	s2	s5	Sowing time
	+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					+-----+-----+-----+-----+-----+					

# History

## Motivation: Sugar beets - A split-plot experiment

```
data(beets, package='pbkrtest')
head(beets)

##   harvest  block  sow yield sugpct
## 1   harv1 block1 sow3 128.0   17.1
## 2   harv1 block1 sow4 118.0   16.9
## 3   harv1 block1 sow5  95.0   16.6
## 4   harv1 block1 sow2 131.0   17.0
## 5   harv1 block1 sow1 136.5   17.0
## 6   harv2 block2 sow3 136.5   17.0

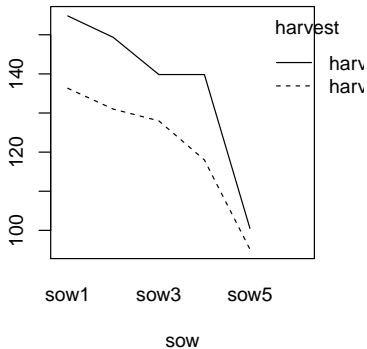
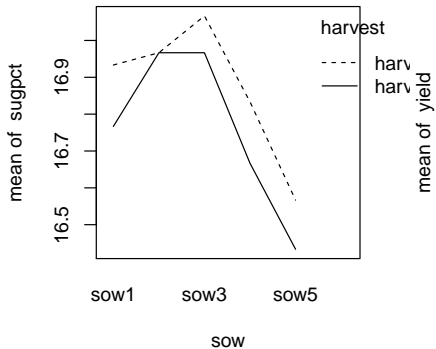
library(doBy)
library(lme4)
```



# History

## Motivation: Sugar beets - A split-plot experiment

```
par(mfrow=c(1,2))  
with(beets, interaction.plot(sow, harvest, sugpkt))  
with(beets, interaction.plot(sow, harvest, yield))
```



# History

## Motivation: Sugar beets - A split-plot experiment

- ▶ For simplicity we assume that there is no interaction between sowing and harvesting times.
- ▶ A typical model for such an experiment would be:

$$y_{hbs} = \mu + \alpha_h + \beta_b + \gamma_s + U_{hb} + \epsilon_{hbs}, \quad (1)$$

where  $U_{hb} \sim N(0, \omega^2)$  and  $\epsilon_{hbs} \sim N(0, \sigma^2)$ .

- ▶ Notice that  $U_{hb}$  describes the random variation between whole-plots (within blocks).

# History

## Motivation: Sugar beets - A split-plot experiment

As the design is balanced we may make F-tests for each of the effects as:

```
beets$bh <- with(beets, interaction(block, harvest))
summary(aov(sugpct ~ block + sow + harvest +
            Error(bh), data=beets))
```

```
##
## Error: bh
##          Df Sum Sq Mean Sq F value Pr(>F)
## block      2  0.0327   0.0163    2.58   0.28
## harvest    1  0.0963   0.0963   15.21   0.06
## Residuals  2  0.0127   0.0063
##
## Error: Within
##          Df Sum Sq Mean Sq F value Pr(>F)
## sow        4    1.01   0.2525   101 5.7e-13
## Residuals 20    0.05   0.0025
```

Notice: the F-statistics are  $F_{1,2}$  for harvest time and  $F_{4,20}$  for sowing time.

# History

## Motivation: Sugar beets - A split-plot experiment

Using `lmer()` from **lme4** we can fit the models and test for no effect of sowing and harvest time as follows:

```
beetLarge <- lmer(sugpct ~ block + sow + harvest +  
                  (1 | block:harvest), data=beets, REML=FALSE)  
beet_no.harv <- update(beetLarge, .~. - harvest)  
beet_no.sow <- update(beetLarge, .~. - sow)
```

# History

## Motivation: Sugar beets - A split-plot experiment

The LRT based  $p$ -values are anti-conservative: the effect of harvest appears stronger than it is.

```
anova(beetLarge, beet_no.sow) %>% as.data.frame
```

##		Df	AIC	BIC	logLik	deviance	Chisq	Chi	Df	Pr(>Chisq)
##	beet_no.sow	6	-2.795	5.612	7.398	-14.8	NA	NA		NA
##	beetLarge	10	-79.998	-65.986	49.999	-100.0	85.2		4	1.374e-17

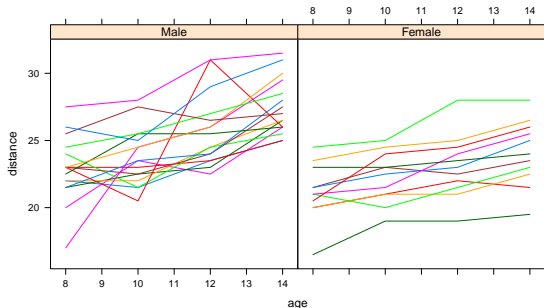
```
anova(beetLarge, beet_no.harv) %>% as.data.frame
```

##		Df	AIC	BIC	logLik	deviance	Chisq	Chi	Df	Pr(>Chisq)
##	beet_no.harv	9	-69.08	-56.47	43.54	-87.08	NA	NA		NA
##	beetLarge	10	-80.00	-65.99	50.00	-100.00	12.91		1	0.0003261

# History

## Motivation: A random regression problem

The change with age of the distance between two cranial distances was observed for 16 boys and 11 girls from age 8 until age 14.



# History

## Motivation: A random regression problem

Plot suggests:

$$dist_{[i]} = \alpha_{sex[i]} + \beta_{sex[i]} age_{[i]} + A_{Subj[i]} + B_{Subj[i]} age_{[i]} + e_{[i]}$$

with  $(A, B) \sim N(0, \mathbf{S})$ .

ML-test of  $\beta_{boy} = \beta_{girl}$ :

```
ort1ML<- lmer(distance ~ age + Sex + age:Sex + (1 + age | Subject),  
              REML = FALSE, data=Orthodont)  
ort2ML<- update(ort1ML, .~. - age:Sex)  
as.data.frame(anova(ort1ML, ort2ML))
```

##		Df	AIC	BIC	logLik	deviance	Chisq	Chi	Df	Pr(>Chisq)
##	ort2ML	7	446.8	465.6	-216.4	432.8	NA	NA		NA
##	ort1ML	8	443.8	465.3	-213.9	427.8	5.029		1	0.02492

# History

## Our goal

Our goal is to extend the tests provided by `lmer()`.

There are two issues here:

- ▶ The choice of test statistic and
- ▶ The reference distribution in which the test statistic is evaluated.

Implement Kenward-Roger approximation.

Implement parametric bootstrap.

Implement Satterthwaite approximation (not yet released)



# The Kenward–Roger approach

## The Kenward–Roger modification of the $F$ -statistic

For multivariate normal data

$$Y_{n \times 1} \sim N(\mathbf{X}_{n \times p} \boldsymbol{\beta}_{p \times 1}, \boldsymbol{\Sigma})$$

we consider the test of the hypothesis

$$\mathbf{L}_{d \times p} \boldsymbol{\beta} = \boldsymbol{\beta}_0$$

where  $\mathbf{L}$  is a regular matrix of estimable functions of  $\boldsymbol{\beta}$ .

With  $\hat{\boldsymbol{\beta}} \sim N_d(\boldsymbol{\beta}, \boldsymbol{\Phi})$ , a Wald statistic for testing  $\mathbf{L}\boldsymbol{\beta} = \boldsymbol{\beta}_0$  is

$$W = [\mathbf{L}(\hat{\boldsymbol{\beta}} - \boldsymbol{\beta}_0)]^\top [\mathbf{L}\boldsymbol{\Phi}\mathbf{L}^\top]^{-1} [\mathbf{L}(\hat{\boldsymbol{\beta}} - \boldsymbol{\beta}_0)]$$

which is asymptotically  $W \sim \chi_d^2$  under the null hypothesis.

# The Kenward–Roger approach

The Kenward–Roger modification of the  $F$ -statistic

A scaled version of  $W$  is

$$F = \frac{1}{d}W$$

which is asymptotically  $F \sim \frac{1}{d}\chi_d^2$  under the null hypothesis – which we can think of as the limiting distribution of an  $F_{d,m}$ -distribution as  $m \rightarrow \infty$

To account for the fact that  $\Phi$  is estimated from data, we must come up with a better estimate of the denominator degrees of freedom  $m$  (better than  $m = \infty$ ).

That was what Kenward and Roger worked on...

# The Kenward–Roger approach

## The Kenward–Roger modification of the $F$ -statistic

The linear hypothesis  $\mathbf{L}\boldsymbol{\beta} = \boldsymbol{\beta}_0$  can be tested via the Wald-type statistic

$$F = \frac{1}{r}(\hat{\boldsymbol{\beta}} - \boldsymbol{\beta}_0)^\top \mathbf{L}^\top (\mathbf{L}^\top \boldsymbol{\Phi}(\hat{\boldsymbol{\sigma}}) \mathbf{L})^{-1} \mathbf{L}(\hat{\boldsymbol{\beta}} - \boldsymbol{\beta}_0)$$

- ▶  $\boldsymbol{\Phi}(\boldsymbol{\sigma}) = (\mathbf{X}^\top \boldsymbol{\Sigma}(\boldsymbol{\sigma}) \mathbf{X})^{-1} \approx \text{Cov}(\hat{\boldsymbol{\beta}})$ ,  $\hat{\boldsymbol{\beta}}$  REML estimate of  $\boldsymbol{\beta}$
- ▶  $\hat{\boldsymbol{\sigma}}$ : vector of REML estimates of the elements of  $\boldsymbol{\Sigma}$

# The Kenward–Roger approach

## The Kenward–Roger modification of the $F$ -statistic

Kenward and Roger (1997) modify the test statistic

- ▶  $\Phi$  is replaced by an improved small sample approximation  $\Phi_A$

Furthermore

- ▶ the statistic  $F$  is scaled by a factor  $\lambda$ ,
- ▶ denominator degrees of freedom  $m$  are determined

such that the approximate expectation and variance are those of a  $F_{d,m}$  distribution.

# The Kenward–Roger approach

The Kenward–Roger modification of the  $F$ -statistic

- ▶ Consider only situations where

$$\Sigma = \sum_i \sigma_i \mathbf{G}_i, \quad \mathbf{G}_i \text{ known matrices}$$

- ▶ Variance component and random coefficient models satisfy this restriction.
- ▶  $\Phi_A(\hat{\sigma})$  depends now only on the first partial derivatives of  $\Sigma^{-1}$ :

$$\frac{\partial \Sigma^{-1}}{\partial \sigma_i} = -\Sigma^{-1} \frac{\partial \Sigma}{\partial \sigma_i} \Sigma^{-1}.$$

- ▶  $\Phi_A(\hat{\sigma})$  depends also on  $\text{Var}(\hat{\sigma})$ .
- ▶ Kenward and Roger propose to estimate  $\text{Var}(\hat{\sigma})$  via the inverse expected information matrix.

# The Kenward–Roger approach

## The Kenward–Roger modification of the $F$ -statistic

### The modification of the $F$ -statistic by Kenward and Roger

- ▶ yields the exact  $F$ -statistic for balanced mixed classification nested models or balanced split plot models (Alnosaier, 2007).
- ▶ Simulation studies (e.g. Spilke, J. et al.(2003)) indicate that the Kenward-Roger approach perform mostly better than alternatives (like Satterthwaite or containment method) for blocked experiments even with missing data.

# The Kenward–Roger approach

The Kenward–Roger modification of the  $F$ -statistic

**lme4** (Bates, D., Maechler, M, Bolker, B., Walker, S. 2014) provides efficient estimation of linear mixed models.

**lme4** provides most matrices and estimates needed to implement a Kenward-Roger approach.

**pbkrtest** (Halekoh, U., Højsgaard, S., 2014) provides a “straight forward” transcription of the description in the article of Kenward and Roger, 1997.

# The Kenward–Roger approach

## The Kenward–Roger modification of the $F$ -statistic

The Kenward–Roger approach yields the same results as the anova-test:

```
beetLarge <- update(beetLarge, REML=TRUE)
beet_no.harv <- update(beet_no.harv, REML=TRUE)
```

Test for harvest effect:

```
KRmodcomp(beetLarge, beet_no.harv)

## F-test with Kenward-Roger approximation; computing time: 0.06 sec.
## large : sugpct ~ block + sow + harvest + (1 | block:harvest)
## small : sugpct ~ block + sow + (1 | block:harvest)
##      stat   ndf   ddf F.scaling p.value
## Ftest 15.2   1.0   2.0       1     0.06
```



# The Kenward–Roger approach

The Kenward–Roger modification of the  $F$ -statistic

For the cranial distances data the Kenward and Roger modified  $F$ -test yields

```
formula(ort1ML)

## distance ~ age + Sex + age:Sex + (1 + age | Subject)

formula(ort2ML)

## distance ~ age + Sex + (1 + age | Subject)

ort1<- update(ort1ML, .~., REML = TRUE)
ort2<- update(ort2ML, .~., REML = TRUE)
```

# The Kenward–Roger approach

The Kenward–Roger modification of the  $F$ -statistic

```
KRmodcomp(ort1, ort2)

## F-test with Kenward-Roger approximation; computing time: 0.11 sec.
## large : distance ~ age + Sex + (1 + age | Subject) + age:Sex
## small : distance ~ age + Sex + (1 + age | Subject)
##      stat    ndf    ddf F.scaling p.value
## Ftest  5.12   1.00 25.52         1   0.032
```

The p-value from the  $\chi^2$ -test was 0.0249.

# The Kenward–Roger approach

## Shortcomings of Kenward-Roger

- ▶ The Kenward–Roger approach is no panacea.
- ▶ In the computations of the degrees of freedom we need to compute

$$G_j \Sigma^{-1} G_j$$

where  $\Sigma = \sum_i \sigma_i \mathbf{G}_i$ . Can be space and time consuming!

- ▶ An alternative is a Satterthwaite–kind approximation which is faster to compute. Will come out in next release of **pbkrtest** (code not tested yet). Way faster...
- ▶ What to do with generalized linear mixed models – or even with generalized linear models.
- ▶ **pbkrtest** also provides the parametric bootstrap  $p$ -value. Computationally somewhat demanding, but can be parallelized.

# Parametric bootstrap

We have two competing models; a large model  $f_1(y; \theta)$  and a null model  $f_0(y; \theta_0)$ ; the null model is a submodel of the large model.

```
lg <- update(beetLarge, REML=FALSE)
sm <- update(beet_no.harv, REML=FALSE)
t.obs <- 2*(logLik(lg)-logLik(sm))
t.obs
```

```
## 'log Lik.' 12.91 (df=10)
```

Idea is simple: Draw  $B$  parametric bootstrap samples  $t^1, \dots, t^B$  under the fitted null model  $\hat{\theta}_0$ .

That is; simulate  $B$  datasets from the fitted null model; fit the large and the null model to each of these datasets; calculate the LR-test statistic for each simulated data:

# Parametric bootstrap

```
set.seed(121315)
t.sim <- PBrefdist(lg, sm, nsim=500)
```

The  $p$ -value is the fraction of simulated test statistics that are larger or equal to the observed one:

```
head(t.sim)

## [1] 3.1363 0.6829 0.1203 1.1063 6.8241 7.3922

sum( t.sim >= t.obs ) / length( t.sim )

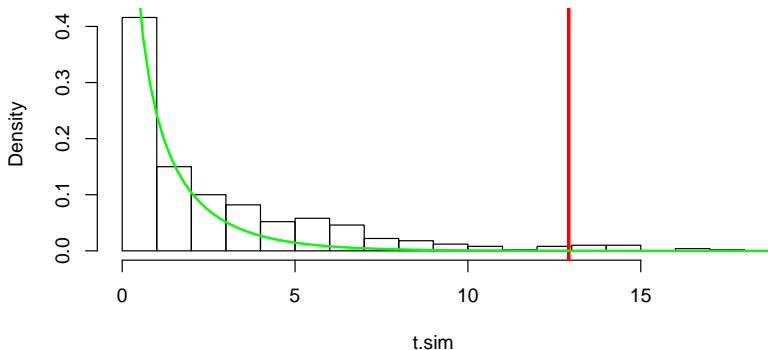
## [1] 0.026
```

# Parametric bootstrap

Interesting to overlay limiting  $\chi^2_1$  distribution and simulated reference distribution:

```
hist(t.sim, breaks=20, prob=T)
abline(v=t.obs, col="red", lwd=3)
f <- function(x){dchisq(x, df=1)}
curve(f, 0, 20, add=TRUE, col="green", lwd=2)
```

Histogram of t.sim



# Parametric bootstrap

Do the same for sowing time:

```
lg <- update(beetLarge, REML=FALSE)
sm <- update(beet_no.sow, REML=FALSE)
t.obs <- 2*(logLik(lg)-logLik(sm))
t.obs
```

```
## 'log Lik.' 85.2 (df=10)
```

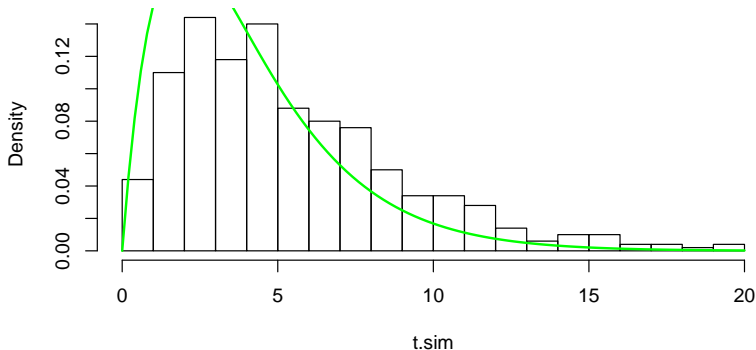
```
set.seed(121315)
t.sim <- PBrefdist(lg, sm, nsim=500)
```

# Parametric bootstrap

Interesting to overlay limiting  $\chi^2_1$  distribution and simulated reference distribution:

```
hist(t.sim, breaks=20, prob=T)
abline(v=t.obs, col="red", lwd=3)
f <- function(x){dchisq(x, df=4)}
curve(f, 0, 20, add=TRUE, col="green", lwd=2)
```

Histogram of t.sim





# Parametric bootstrap

This scheme is implemented as:

R

```
set.seed(121315)
pb <- PBmodcomp(beetLarge, beet_no.harv)
pb

## Parametric bootstrap test; time: 19.17 sec; samples: 1000 extremes: 40;
## large : sugpct ~ block + sow + harvest + (1 | block:harvest)
## small : sugpct ~ block + sow + (1 | block:harvest)
##      stat df p.value
## LRT    11.8  1 0.00059
## PBtest 11.8   0.04096
```

# Parametric bootstrap

In addition we can get  $p$ -values

1. directly via the proportion of sampled  $t_i$  exceeding  $t_{obs}$ ,
2. approximating the distribution of the scaled statistic  $\frac{f}{\bar{t}} \cdot T$  by a  $\chi_f^2$  distribution (Bartlett type correction)  
( $\bar{t}$  is the sample average and  $f$  the difference in the number of parameters between the null and the alternative model)
3. approximating the bootstrap distribution by a  $\Gamma(\alpha, \beta)$  distribution which mean and variance match the moments of the bootstrap sample.
4. approximating the bootstrap distribution by a  $F_{d,m}$  distribution which mean is based on matching mean of the bootstrap sample.

# Parametric bootstrap

```
summary(pb)
```

```
## Parametric bootstrap test; time: 19.17 sec; samples: 1000 extremes: 40;  
## large : sugpct ~ block + sow + harvest + (1 | block:harvest)  
## small : sugpct ~ block + sow + (1 | block:harvest)  
##          stat      df  ddf p.value  
## PBtest    11.82          0.04096  
## Gamma     11.82          0.03510  
## Bartlett   4.05  1.00      0.04416  
## F          11.82  1.00  3.04  0.04042  
## LRT        11.82  1.00      0.00059
```

# Parametric bootstrap

## Parallel computations

Parametric bootstrap is computationally demanding, but multiple cores can be exploited:

```
library(parallel)
nc <- detectCores()
nc
```

```
## [1] 4
```

```
clus <- makeCluster(rep("localhost", nc))
```

# Parametric bootstrap

## Parallel computations

### R

```
set.seed(121315)
pb1 <- PBmodcomp(beetLarge, beet_no.harv)
pb1

## Parametric bootstrap test; time: 19.12 sec; samples: 1000 extremes: 40;
## large : sugpct ~ block + sow + harvest + (1 | block:harvest)
## small : sugpct ~ block + sow + (1 | block:harvest)
##          stat df p.value
## LRT      11.8  1 0.00059
## PBtest 11.8    0.04096

pb2 <- PBmodcomp(beetLarge, beet_no.harv, cl=clus)
pb2

## Parametric bootstrap test; time: 10.00 sec; samples: 1000 extremes: 42;
## large : sugpct ~ block + sow + harvest + (1 | block:harvest)
## small : sugpct ~ block + sow + (1 | block:harvest)
##          stat df p.value
## LRT      11.8  1 0.00059
## PBtest 11.8    0.04296
```

# Parametric bootstrap

## Parallel computations

Results from sugar beets:

**Table:** p-values ( $\times 100$ ) for removing the harvest or sow effect.

	LRT	KR	ParmBoot	Bartlett	Gamma
harvest	0.03	6	4.1	8.3	4.9
sow	<0.001	<0.001	<0.001	<0.001	<0.001

Results for cranial distance data:

**Table:** p-values ( $\times 100$ ) testing  $\beta_{boy} = \beta_{girl}$ .

LRT	KR	ParmBoot	Bartlett	Gamma
2.5	3.3	4.2	4.0	4.2

# Parametric bootstrap

## Parallel computations

The above approaches are computationally intensive but there are possibilities for speedups:

Instead of simulating a fixed number of values  $t^1, \dots, t^M$  for determining the reference distribution used for finding  $p^{PB}$  we may instead introduce a stopping rule saying *simulate until we have found, say 20 values  $t^j$  larger than  $t_{obs}$* . If  $J$  simulations are made then the reported  $p$ -value is  $20/J$ .

Estimating tail-probabilities will require more samples than estimating the mean (and variance) of the reference distribution. Therefore the Bartlett and gamma approaches will require fewer simulations than needed for finding  $p^{PB}$ .

The simulation of the reference distribution can be parallelized onto different processors.

# Small simulation study: A random regression problem

We consider the simulation from a simple random coefficient model (cf. Kenward and Roger (1997, table 4)):

$$y_{it} = \beta_0 + \beta_1 \cdot t_i + A_i + B_i \cdot t_i + \epsilon_{it}$$

with  $\text{cov}(A_i, B_i) = \begin{bmatrix} 0.250 & -0.133 \\ -0.133 & 0.250 \end{bmatrix}$  and  $\text{var}(\epsilon_{it}) = 0.25$ .

There are observed  $i = 1, \dots, 24$  subjects divided in groups of 8. For each group observations are at the non overlapping times  $t = 0, 1, 2$ ;  $t = 3, 4, 5$  and  $t = 6, 7, 8$ .



# Small simulation study: A random regression problem

**Table:** Observed test sizes ( $\times 100$ ) for  $H_0 : \beta_k = 0$  for random coefficient model.

	LR	Wald	ParmBoot	Bartlett	Gamma	KR(R)	KR(SAS)
$\beta_0$	6.8	4.6	5.2	5.2	5.4	4.0	5.4
$\beta_1$	7.3	5.3	6.0	6.0	5.9	5.4	6.3

# Final remarks

- ▶ The functions `KRmodcomp()` and `PBmodcomp()` described here are available in the `pbkrtest` package.
- ▶ The Kenward–Roger approach requires fitting by REML; the parametric bootstrap approaches requires fitting by ML.
- ▶ The required fitting scheme is set by the relevant functions, so the user needs not worry about this.
- ▶ Parametric bootstrap is parallelized using the `snow` package.

## Final remarks

- ▶ Halekoh, U., Højsgaard, S. (2014) *A Kenward-Roger Approximation and Parametric Bootstrap Methods for Tests in Linear Mixed Models The R Package pbkrtest*
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